Neuron

Circuit Mechanisms Underlying Motor Memory Formation in the Cerebellum

Highlights

- Pauses in PN firing drive cerebellar nuclei neurons and rapid forelimb movements
- Pairing PN excitation or inhibition with sensory stimuli causes motor learning
- Movements trained by PN excitation are more accurately timed
- PN excitation and inhibition trigger plasticity at distinct sites in the cerebellum

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In Brief

Lee et al. implicate inhibition of PNs as an instructive signal in motor learning by demonstrating that reductions in cerebellar PN firing are sufficient to both elicit movement and, when paired with sensory stimuli, drive associative motor memory formation.



Article

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SUMMARY

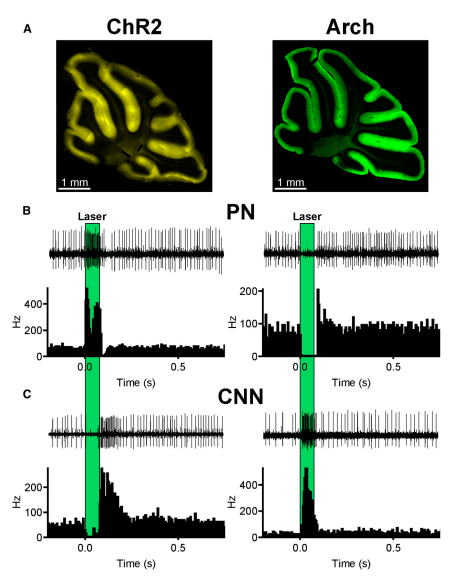
The cerebellum stores associative motor memories essential for properly timed movement; however, the mechanisms by which these memories form and are acted upon remain unclear. To determine how cerebellar activity relates to movement and motor learning, we used optogenetics to manipulate spontaneously firing Purkinje neurons (PNs) in mouse simplex lobe. Using high-speed videography and motion tracking, we found that altering PN activity produced rapid forelimb movement. PN inhibition drove movements time-locked to stimulus onset, whereas PN excitation drove delayed movements time-locked to stimulus offset. Pairing either PN inhibition or excitation with sensory stimuli triggered the formation of robust, associative motor memories; however, PN excitation led to learned movements whose timing more closely matched training intervals. These findings implicate inhibition of PNs as a teaching signal, consistent with a model whereby learning leads first to reductions in PN firing that subsequently instruct circuit changes in the cerebellar nucleus.

INTRODUCTION

To ensure coordination, the brain must make accurate predictions about how to direct movement (Medina, 2011). These predictions are constructed through a process of error-driven learning, and then stored as associative memories in the cerebellum (Albus, 1971; Marr, 1969; Medina, 2011; Raymond et al., 1996). Correlates of such memories have been observed in the firing patterns of cerebellar Purkinje neurons (PNs) as sensory-evoked reductions in the firing rate of PNs in advance of a learned movement (Jirenhed et al., 2007; Lisberger et al., 1994a; Medina and Lisberger, 2008). PNs show high rates of spontaneous activity causing them to powerfully inhibit cerebellar nuclear and vestibular neurons (CNNs), which function as premotor neurons (Person and Raman, 2012). Thus, reductions in spontaneous PN firing in response to the predictive sensory stimulus could, in principle, drive learned movements. Although this disinhibition hypothesis was first suggested four decades ago (Albus, 1971), it is still actively debated (De Zeeuw et al., 2011; Heck et al., 2013; Ito, 1984; Medina, 2011), in part because PN firing patterns do not straightforwardly encode aspects of movement (Cao et al., 2012; Catz et al., 2008; Greger et al., 2004; Kojima et al., 2010; Popa et al., 2013). Moreover, evidence causally linking specific patterns of PN activity to discrete movements has been lacking until very recently (Heiney et al., 2014).

Learning-related changes in PN firing are a hallmark of associative, cerebellum-dependent motor learning, and stimulusinduced reductions in PN firing are proposed to play a key role in learning involving increases in movement (Hesslow and Ivarsson, 1994; Jirenhed et al., 2007; Lisberger et al., 1994a; Medina and Lisberger, 2008). Learning is also hypothesized to involve plasticity downstream of PNs within the cerebellar nuclei (Lisberger et al., 1994b; Miles and Lisberger, 1981; Ohyama and Mauk, 2001; Ohyama et al., 2006; Perrett et al., 1993). Evidence from a variety of learning models suggests that associative memory formation occurs in two stages, with circuit changes occurring first in the cerebellar cortex, followed by later changes in the cerebellar nuclei (Cooke et al., 2004; Kassardjian et al., 2005; Ohyama and Mauk, 2001; Okamoto et al., 2011; Shutoh et al., 2006; Titley et al., 2007). In this scenario, learning-related reductions in PN firing could serve to instruct changes in the cerebellar nuclei, leading, for example, to strengthening of excitatory inputs to CNNs (Maiz et al., 2012; Mauk, 1997; Mauk and Donegan, 1997; Medina, 2011; Medina and Lisberger, 2008; Ohyama and Mauk, 2001; Otis et al., 2012). At a cellular level, disinhibition of CNNs is known to induce activity-dependent forms of long-term potentiation that could support this type of learning (Pugh and Raman, 2006). Also consistent with this idea, genetic deletion of GABAA receptors in PNs results in a memory consolidation defect (Wulff et al., 2009), supporting the notion that inhibition-induced pauses in PN firing are required for the formation of cerebellar memories.

In order to determine how PN firing relates to movement and to explore whether certain patterns of PN activity could drive the formation of associative motor memories, we developed a behavioral paradigm allowing direct manipulation of PN firing



and precise measurement of forelimb movements. We find that inhibiting PNs drives short-latency forelimb movements, while exciting PNs results in forelimb movements delayed to the offset of excitation, results similar to those described recently for postural movements (Witter et al., 2013). Optrode experiments indicate that in both circumstances movement is linked to pauses in PN firing and bursts in downstream CNNs. By pairing PN activity with auditory tones, we demonstrate robust associative learning leading to tone-evoked, predictive forelimb movements. Such learning is accompanied by extensive structural plasticity in the cerebellar nuclei. As suggested by a two-stage model, pairing with either excitation or inhibition of PNs drives learning, but the timing of learned movements trained by PN inhibition is less predictive of the training interval, consistent with observations that plasticity in the cerebellar cortex is required for well-timed, learned movements. These findings demonstrate that reductions in PN firing are sufficient to elicit discrete movements, and that repeatedly pairing such reductions with sensory

(A) Photomicrographs of parasagittal cerebellar sections showing expression of ChR2-eYFP (left) and Arch-GFP (right) selectively in PNs.

(B) In vivo, single-unit optrode recordings from PNs in response to 75-ms light activation of ChR2 (left) or Arch (right) in awake mice. Laser pulse duration is indicated by green boxes. Single extracellular traces are shown above the peri-stimulus time histograms for each cell.

(C) In vivo recordings from CNNs in ChR2 (left) and Arch (right) mice in response to 75-ms light pulses delivered to PN axons in the cerebellar nucleus. See also Figures S2 and S3.

stimuli leads to the formation of associative motor memories, implicating inhibition of PNs as an instructive mechanism for driving learning-related changes in the cerebellar nuclei.

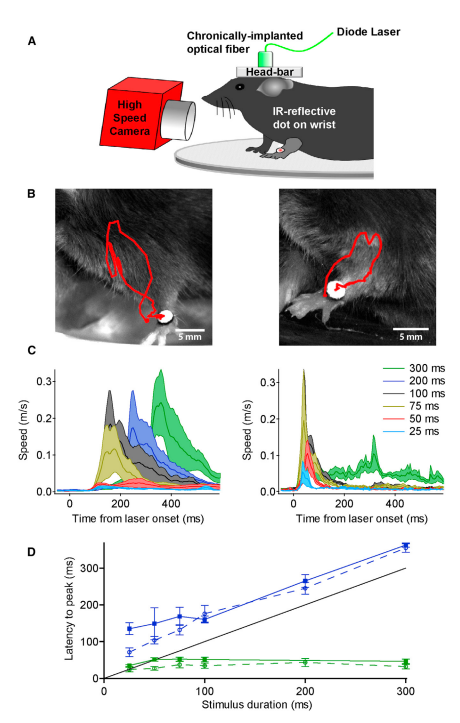
RESULTS

To examine the underlying mechanisms by which PNs contribute to movement and associative motor memory formation, we exploited Cre-conditional transgenic mice to direct expression of the excitatory opsin channelrhodopsin-2 fused to eYFP (ChR2) or the inhibitory opsin archaerhodopsin-3 fused to eGFP (Arch) in PNs. Expression of ChR2 or Arch was selective for PNs throughout the cerebellum and was apparent in axons projecting to CNNs (Figures 1A and S1). Electrophysiological experiments in brain slices showed that ChR2 activation generates large inward currents and rapid increases in PN firing followed by brief pauses in

spontaneous firing activity, while Arch activation generates large outward currents and pauses in the spontaneous firing of PNs (Figure S2). Light-evoked responses and fluorescence were absent from other types of neurons in the cerebellar cortex such as molecular layer interneurons, granule cells, and Golgi cells.

Inhibition of Spontaneously Firing PNs Drives Movement

To examine behavioral consequences of transient modulation of PN activity, we developed an awake, head-fixed mouse preparation amenable to simultaneous opsin excitation and extracellular recording. Activation of ChR2 for 75 ms led to robust increases in the simple spike (SSp) firing frequency of PNs (Figure 1B, left, peak frequency = 363.4 ± 40.7 Hz, n = 11), and in most PNs the increase was followed by pauses in SSps occurring upon cessation of laser illumination (pause duration 35.0 ± 4 ms in n = 9/11 PNs). Recordings of downstream CNN firing showed that ChR2-mediated activation of PNs strongly inhibits CNNs during laser illumination, and that large increases in firing rate occur



upon cessation of the laser pulse (Figure 1C, left, peak frequency = 275.3 ± 40.6 Hz, n = 18). In contrast to the ChR2 results, inhibiting PNs via Arch rapidly silenced PN SSp activity (Figure 1B, right, pause duration 99.1 \pm 7.4 ms, n = 16) and strongly excited CNNs (Figure 1C, right, peak frequency = 219.6 ± 42.4 Hz, n = 15) during laser illumination. Statistical comparison of peak ChR2-driven rebound firing and peak Arch-driven firing in CNNs indicated no significant differences (p = 0.33, two-tailed, paired t test). Systematically varying pulse

Figure 2. Optogenetically Elicited, Rapid Forelimb Movements

(A) Illustration of head-fixed animal preparation used for behavioral studies. A high-speed video camera coupled with infrared (IR) illumination allowed movement of the wrist, tagged with an IRreflective dot, to be tracked offline using motiontracking software.

(B) Shown is a single frame taken from the end of a video in which a 100-ms laser pulse was delivered through a chronically implanted fiber optic in the forelimb region of the ipsilateral cerebellar cortex. The resulting ChR2 (left) and Arch (right) forelimb trajectories are traced in red.

(C) Mean forelimb speed versus time relative to the onset of laser pulses of the indicated durations. Note that in ChR2 mice (left, n = 4) movement onset is time-locked to the end of laser illumination, while in Arch mice (right, n = 4) movement occurs independently of pulse duration with a fixed delay following illumination onset.

(D) Delay to peak movement speed (solid lines) and peak CNN firing frequency (dotted lines) are plotted together as a function of laser pulse duration. Data are plotted for ChR2 (blue, n = 4 animals/ 7 cells) and Arch (green, n = 4 animals/9 cells) mice. Note that in ChR2 mice movement and peak firing occur with a fixed delay from the end of the laser pulses (indicated by black line), whereas in Arch mice movement and peak CNN firing occur at a fixed delay from pulse onset. See also Figures S2 and S3 and Movies S1 and S2.

duration demonstrated that Arch stimulation led to CNN firing that was time locked to the onset of illumination, while ChR2 stimulation led to CNN inhibition followed by excitation upon offset of laser illumination (Figure S3). Taken together, these findings indicate that the firing frequencies of PNs and CNNs are robustly and differentially modulated by optogenetic stimuli, and that briefly inhibiting PNs is sufficient to produce high-frequency firing of CNNs.

To test whether manipulation of PN/ CNN excitability generates motor output we stereotaxically implanted optical fibers in the forelimb motor region of the anterior cerebellar lobe (Figure S1). Movement of the forelimb was monitored in head-fixed mice and measured using high-speed

videography (Chettih et al., 2011; Heiney et al., 2014) and motion tracking analysis (Figure 2; Movie S1). Inhibiting PNs via activation of Arch led to rapid and stereotyped upward forelimb movements during laser illumination that reached peak velocities of 0.3–1.5 m/s (Figures 2 and S4). By comparison, excitation of PNs via ChR2 led to delayed movement of similar speed and magnitude, but with onsets time locked to the laser pulse termination (Figures 2 and S4). This relationship between stimulus onset/offset and movement onset can be visually compared by

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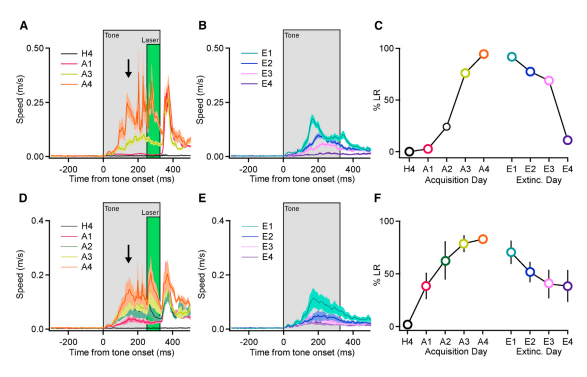


Figure 3. Pairing ChR2 Activation in PNs with Auditory Tones Leads to Predictive, Tone-Evoked Forelimb Movements

(A) Mean speed (± SEM indicated by lighter shading) of forelimb movement for a single animal across all trials on indicated days of training (H4, habituation day 4; A1, 3, and 4, acquisition training days 1, 3, and 4). All analyzed trials are included. The timing of the tone (325 ms) and laser (75 ms) pulses are indicated with gray and green boxes, respectively, and the arrow indicates the LR. Note the gradual appearance of tone-evoked movement with acquisition training and abolishment of this movement with extinction training.

(B) Mean movement speed during extinction training (E1-4, extinction days 1-4) for the same animal.

(C) Summary plot for this animal indicating the percent of LRs detected on each acquisition and extinction training. Colored symbols correspond to mean speed-versus-time traces in (A) and (B).

(D and E) Group average of mean speed profiles during acquisition (D) and extinction (E), n = 4 mice for each.

(F) Summary of percent LR ± SEM (n = 4 mice) across days. See also Figures S5 and S6 and Movie S3.

viewing Movie S2. Across animals, the average timing relative to laser illumination of peak CNN firing and peak movement speed in the two mouse lines indicates that by driving CNN firing, PN pauses lead to the observed forelimb movements (Figure 2D).

Robust Motor Memories Are Induced by Pairing PN Activation with Auditory Tones

Cerebellum-dependent, associative motor learning has been hypothesized to involve forms of synaptic plasticity triggered by teaching signals from olivo-cerebellar climbing fibers at multiple sites within the cerebellar circuit (Maiz et al., 2012; Medina and Lisberger, 2008; Raymond et al., 1996), yet the mechanisms underlying this learning are under active debate (Ke et al., 2009; Medina, 2011; Schonewille et al., 2011). Since both climbing fiber-dependent and -independent PN plasticity rely on PN depolarization, we tested whether synchronous PN depolarization can serve as a teaching signal. ChR2 mice were trained by pairing 2-kHz tones with pulses of laser illumination (75-100 ms) delivered 250 ms after tone onset. Each day, training consisted of 90 tone/laser pairings and 10 interleaved tone-alone trials extended over 4 days of acquisition training. Following this acquisition training, mice were subjected to extinction training in which they were presented 100 tone-alone trials on each day. Within 2-3 days, acquisition training led to robust and statistically significant (one-way ANOVA, F(2,8) = 4.403, p = 0.00168) learned responses (LRs) evident as tone-evoked forelimb movements (Figures 3, S5, and S6; Movie S3), which occurred in 83% ± 4.4% of trials and were rapidly extinguished over 3-4 days of extinction training (post hoc Tukey's HSD test, pairwise comparison to H4, E3 p = 0.143, E4 p = 0.202). Kinematic analyses showed that these learned movements involved the same limb and were similar in direction and temporal profile to those evoked by the laser pulses (Figures S5 and S6), but occurred earlier in time, as expected for predictive cerebellar learning (Medina, 2011; Ohyama et al., 2003). These findings indicate that synchronous PN depolarization can trigger the formation of associative motor memories allowing sensory stimuli to drive predictive forelimb movements.

Pairing PN Inhibition with Auditory Tones Also Leads to Associative Learning

Evidence suggests that cerebellum-dependent associative learning involves circuit changes not only in the cerebellar cortex, but also in the cerebellar nuclei (Broussard and Kassardjian, 2004; Gao et al., 2012; Lisberger, 1994; Miles and Lisberger,

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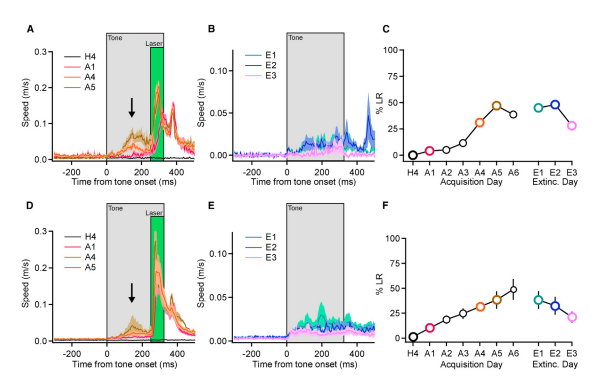


Figure 4. Pairing Arch Activation in PNs with Auditory Tones Leads to Predictive, Tone-Evoked Forelimb Movements

(A) Mean speed (± SEM indicated by lighter shading) of forelimb movement for a single animal across all trials on indicated days of training (H4, habituation day 4; A1, 4, and 5, acquisition training days 1, 4, and 5). All analyzed trials are included. The timing of the tone (325 ms) and laser (75 ms) pulses are indicated with gray and green boxes, respectively, and the arrow indicates the LR. Note the gradual appearance of tone-evoked movement with acquisition training and abolition of this movement with extinction training.

(B) Mean speed during extinction training (E1-3, extinction days 1-3) for the same animal.

(C) Summary plot for this animal indicating the percent of LRs detected on each acquisition and extinction training. Colored symbols correspond to mean speed-versus-time traces in (A) and (B).

(D and E) Group average of mean speed profiles for Arch mice during acquisition (D, n = 6 mice) and extinction (E, n = 3 mice).

(F) Summary of percent LR ± SEM (n = 2–6 mice) across days. See also Figures S5 and S6 and Movie S4.

1981: Ohvama et al., 2006: Raymond et al., 1996). In principle, pauses in PN activity driven by either complex spikes (Maiz et al., 2012) or those developed in response to learning may instruct such changes by conveying transient periods of disinhibition to CNNs leading to the potentiation of tone-driven inputs to those CNNs. Mechanistically, NMDA receptor-dependent forms of long-term potentiation, which have been observed in vitro, could serve as the basis for this learning (Pugh and Raman, 2006). To test whether PN pauses are also sufficient to induce LRs we applied an associative training paradigm similar to that described above for ChR2, but using Arch to synchronously inhibit PNs. Pairing tones with Arch-driven PN pauses instructed tone-evoked forelimb movements (Figures 4, S5, and S6). LRs in Arch mice developed over several days of acquisition training and extinguished in response to tone-alone training (one-way ANOVA, F(2,10) = 4.797, p = 0.0001, post hoc Tukey's HSD test, in pairwise comparison to H4, E3 p = 0.3234); however, the percent of LRs was smaller than that observed for ChR2, and the individual LRs were of smaller amplitude (Table S1). These results indicate that disinhibition of CNNs alone is sufficient to drive associative motor learning, and suggest that synchronous pauses in the firing of groups of PNs can trigger circuit changes in the CNN that underlie associative learning.

PN Inhibition-Induced Learning Leads to Mossy Fiber Sprouting in Cerebellar Nuclei

A recent study demonstrated that paired, but not unpaired, tone/ air-puff training triggered significant structural changes in the cerebellar nuclei involving an increase in the number of mossy fiber endings related to the auditory tone (Boele et al., 2013). These findings support the hypothesis that associative conditioning leads to strengthening of specific mossy fiber to CNN connections, thereby contributing to LRs. We set out to examine whether similar structural plasticity occurs in the cerebellar nuclei of Arch-trained animals. To first assess which areas of the cerebellar nuclei receive inputs from the PNs silenced by Arch activation during training, a small amount of dextran-conjugated Alexa 488 dye was injected into the forelimb motor region of lobulus simplex, the same site as we implanted our optical fibers (compare Figures S1 and S7). Dye-filled axonal projections and terminals were found within discrete regions of the cerebellar nuclei, including large portions of the anterior interpositus nucleus (IntA) and dorsolateral protuberance of the medial nucleus (MedDL) (Figure 5A), regions representing forelimb areas in rats (Pardoe and Apps, 2002). Next, we examined whether the number of mossy fiber terminals increased within these regions of the cerebellar nuclei after associative learning. To

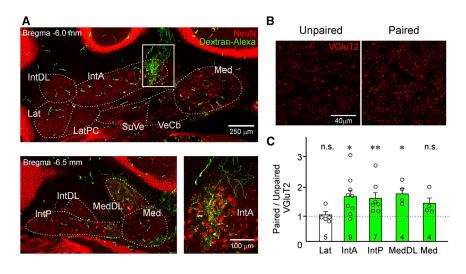


Figure 5. Associative Motor Learning Induces Formation of VGluT2+ Terminals in a Subset of Cerebellar Nuclei in L7-Arch Mice (A) Coronal sections, counterstained with the neuronal marker, NeuN (red), show dextran-conjugated Alexa 488-filled PN axons (thin green lines) that originate from the same site as our optic fiber placements. Dye-filled PN axons are apparent within the anterior interpositus (IntA), posterior interpositus (IntP), medial (Med), and dorsolateral protuberance of the medial nucleus (MedDL). Green objects of large diameter are blood vessels, and not PN axons (see Figure S7). Note that the top- and bottom-left are at different rostro-caudal positions as indicated in the top-left corner of each image. In many cases synaptic terminals can be made out within the individual nuclei (bottom-right, corresponds to white box in top image). Dorsolateral hump of the interpositus (IntDL), lateral (Lat). parvicellular lateral (LatPC), superior vestibular (SuVe), vestibulocerebellar (VeCb).

(B) Confocal micrographs of mossy fiber synaptic terminals labeled with the vesicular glutamate transporter 2 (VGluT2, red) in the MedDL of paired (right) and unpaired (left) mice.

(C) The ratio of the number of VGluT2+ puncta in various cerebellar nuclei of paired versus littermate-matched unpaired controls. Green bars indicate nuclei in which dye-filled PN axons were detected. Data are represented as mean value \pm SEM; open circles indicate individual pairs. *p < 0.05, **p < 0.01, and ns, no statistical difference.

do this we divided sibling cage-mates into two different groups-paired and unpaired. The paired group received 4 days of tone-laser paired training as described in Figure 4. The unpaired group was presented the same number of tone and laser stimuli over 4 days; however, these stimuli were presented in an unpaired, pseudorandomized fashion. To assess structural changes in mossy fiber inputs in the cerebellar nuclei, the number of mossy fiber puncta in the paired mice was compared against that in sibling-matched unpaired mice (Figure 5B). The number of VGluT2-positive puncta was significantly higher in regions of the cerebellar nuclei receiving PN terminals originating from the forelimb area in paired versus unpaired mice (Figure 5C, MedDL, p = 0.026, n = 4 matched nuclear regions; IntA, p = 0.024, n = 9; IntP, p = 0.0023, n = 7; paired t test, data from three or four matched mouse pairs). In comparison, no significant changes in mossy fiber puncta were observed in the lateral nucleus (Lat, p = 0.69, n = 5; paired t test, data from three matched mouse pairs), consistent with the absence of dye-filled PN axons in this region of the cerebellar nucleus (see Experimental Procedures). These results indicate that the LRs observed in Arch-trained mice are accompanied by structural plasticity of mossy fiber inputs in the cerebellar nucleus similar to normal associative learning (Boele et al., 2013).

Learned Movements Trained with PN Excitation versus Inhibition Show Differences in Timing

A critical aspect of associative motor memories is that they are adaptively timed, enabling predictive movements to be executed at precise times during sensory stimuli. Lesion studies suggest that these memories are stored in both the cerebellar cortex and nuclei, but that information related to the timing of learned movements resides in the cerebellar cortex (Ohyama et al., 2003, 2006; Perrett et al., 1993). To compare the timing of learned movements generated by ChR2 versus Arch training, we made kinematic measurements of LRs resulting from two different training intervals. For ChR2-trained animals, despite being similar in amplitude, the time courses of average, peak LRs were different, precisely anticipating the ends of the 250or 500-ms training intervals (Figure 6A; Table S1). In contrast, LRs resulting from 250- and 500-ms training in Arch animals were similarly timed such that the LRs in mice trained with 500-ms intervals showed peak movement speed early in the stimulus (Figure 6B). Comparison between ChR2 and Arch of the latencies to peak movement speeds for the 500-ms training intervals (Figure 6C) further support the conclusion that learned movements are significantly better timed in ChR2trained animals.

The differences in timing of LRs described above suggest that learning that results from PN inhibition occurs downstream within the cerebellar circuit, and a likely mechanism for this learning involves potentiation of tone-related mossy fiber inputs to CNNs (Boele et al., 2013; Miles and Lisberger, 1981; Ohyama et al., 2006; Raymond et al., 1996). Such a mechanism leads to the prediction that movements evoked by PN inhibition would be facilitated if accompanied by a sensory stimulus (e.g., tone) to which learning has occurred. We tested this prediction in Arch mice by comparing movements in response to the tone alone and to those evoked by the same tone plus a laser pulse timed to elicit PN pauses at the peak of the LR (Figure 7A). As shown in Figure 4, tone-alone stimuli evoked no movement prior to learning. Following 4 days of paired training, inhibiting PNs using weak laser pulses facilitated tone-evoked LRs by $361\% \pm 63\%$ (tone-plus-laser compared to tone-alone in interleaved trials, n = 3 mice, see Figure 7), leading to a significantly larger movement compared to the sum of movements evoked by laser stimulation and tone stimulation alone (sum = $202\% \pm 72\%$, p < 0.005, two-tailed, paired t test). These results indicate that the contributions of PN pauses to movement are influenced by

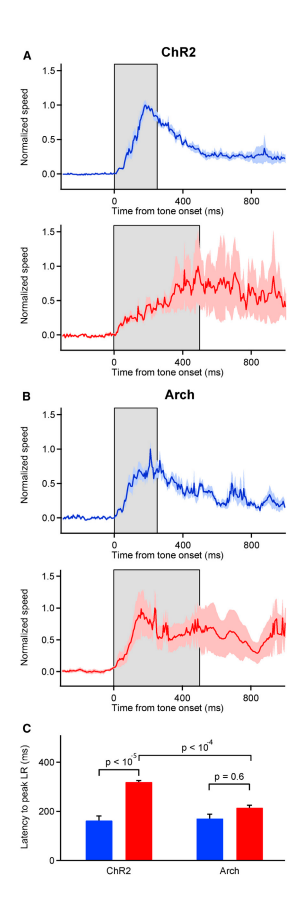


Figure 6. ChR2 Training, but Not Arch Training, Leads to Well-Timed, Learned Movements

(A) Average, normalized movement speed for ChR2 mice in tone-alone LRs occurring between acquisition day 3 and extinction day 1 trained with 250-ms (blue, n = 4 mice) or 500-ms intervals (red, n = 4 mice). Gray boxes indicate interval between tone and laser pulse onset. Lighter shading represents ± SEM across mice.

(B) Average, normalized movement speed in tone-alone LRs for Arch mice trained with 250- (blue, n = 7 mice) or 500-ms (red, n = 5 mice) intervals.

(C) Mean (\pm SEM) of the delay to the peak LR for all trials included in the averages in (A) and (B). LR timing in ChR2-trained mice is significantly different between the 250- and 500-ms intervals and between ChR2 and Arch for the 500-ms interval, but not for Arch-trained mice between the 250- and 500-ms intervals. p values determined using a two-tailed, Mann-Whitney U test. See also Figure S7 and Table S1.

whether or not learning has occurred, and support the hypothesis that inhibition of PN firing can instruct memory-related changes in the cerebellar nuclei, a potential mechanistic substrate for "offline" consolidation from the cerebellar cortex to the cerebellar nuclei.

DISCUSSION

Here we combined optogenetics, high-speed videography, and motion tracking to explore how modulation of the firing of groups of PNs in the simplex lobe acutely affects movement and how coupling changes in PN firing with sensory experience can drive motor memory formation. This novel paradigm allows for the creation of artificial memories, powerfully demonstrating the generality of basic principles of cerebellar associative learning uncovered in classical eyeblink conditioning and vestibuloocular reflex plasticity. Our results also implicate critical functional roles for PN inhibition in the cerebellar circuit. On a fast timescale, reductions in PN activity lead to movement, while on a longer timescale, reductions in PN activity lead to behavioral learning, likely by stimulating increases in mossy fiber input to CNNs. Finally, by engaging distinct circuit elements with ChR2 and Arch training, we show that learned forelimb movements exhibit different timing profiles, providing support for the hypothesis that cerebellar learning results from circuit modifications in both the cerebellar cortex and nucleus with plasticity in the cerebellar cortex required for precisely timed learned movements.

Synchronous Reductions in PN Firing Drive Forelimb Movements

Despite extensive study, direct links between PN activity and specific aspects of movement have been unclear. While electrical microstimulation of the cerebellar nuclei reliably triggers limb movements (Ekerot et al., 1995; Rispal-Padel et al., 1981, 1982; Schultz et al., 1976, 1979), and microstimulation of floccular and vermal regions of cerebellar cortex leads to movements of the head, parts of the face, or eyes (Cohen et al., 1965; De Zeeuw and Koekkoek, 1997; Esakov and Pronichev, 2001; Lisberger, 1994; Noda and Fujikado, 1987; Ron and Robinson, 1973), there are no reports in the literature of limb movements evoked by electrical stimulation of the cerebellar cortex. Interestingly, microstimulation in more lateral areas of cerebellar cortex located

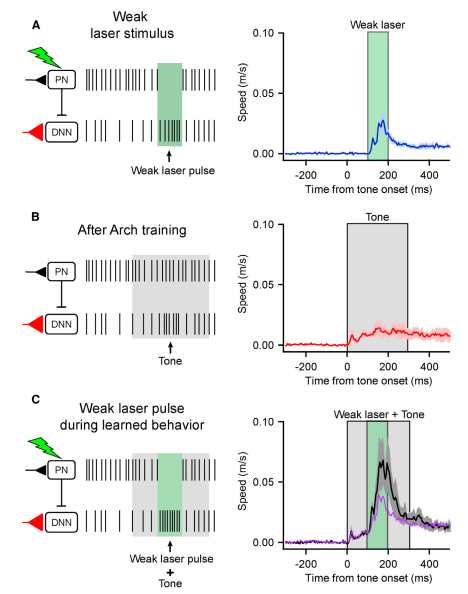


Figure 7. Arch Training Alters the Excitability of CNNs in a Tone-Dependent Manner

Mice were associatively trained by pairing tones and Arch activation as shown in Figure 4. Each panel indicates on the left the hypothesized effects on PN and CNN firing in response to the stimulus and on the right the mean movement speed \pm SEM for three mice.

(A) Mean movements evoked by weak laser pulses.

(B) Mean tone-evoked LRs after 4 days of training.
(C) Mean movements in response to the tone and weak laser pulse delivered simultaneously. Note that the resulting augmented LRs (black) are larger than the arithmetic sum of the laser-evoked plus the learned movements (purple).

Modulation of PN Activity in Conjunction with Sensory Stimuli Induces Motor Memories

We also show that specific PN activity patterns, when paired with sensory stimuli, can drive robust, associative motor learning (Figures 3 and 4). The resulting learned movements closely resemble the acutely evoked movements recorded in our study, consistent with the somatotopic map known to exist in cerebellum (Apps and Hawkes, 2009). The ChR2induced, artificial memories described here also have a number of properties similar to natural forms of cerebellar learning such as eyeblink conditioning and vestibuloocular reflex plasticity (Raymond et al., 1996). These include the rate at which learning and extinction occur, the predictive timing of LRs, the registration between training interval and learned movement timing, and the structural plasticity observed in the nucleus.

near those under study here has been reported to suppress learned eyeblink movements (Hesslow, 1994).

Given the well-established correlation between reductions in PN firing and learned movements, the issue of how PN firing contributes to movement is critical (De Zeeuw et al., 2011; Jirenhed et al., 2007; Lisberger et al., 1994a; Medina and Lisberger, 2008). Our findings comparing the effects of stimulating ChR2 and Arch in PNs indicate that synchronous pauses in spontaneous firing of PNs in the simplex lobe are sufficient to elicit rapid forelimb movements (Figures 1 and 2). These results are in line with recent reports showing facial movements in response to ChR2-elicited synaptic inhibition of PNs (Heiney et al., 2014) and delayed postural movements elicited by ChR2 stimulation of PNs (Witter et al., 2013). Taken together, the findings imply that populations of functionally related PNs promote movement by synchronizing the inter-spike intervals on a fast timescale thereby disinhibiting CNNs to drive movements (Person and Raman, 2012). The findings suggest a conserved mechanism by which cerebellar circuits can make use of sensorimotor information to prompt adaptive movements, at least within the intermediate cerebellum.

The learning in response to Arch stimulation demonstrates that PN inhibition can trigger the formation of associative motor memories. A likely mechanism for this learning involves disinhibition of CNNs leading to an enhancement in the strength of mossy fiber collateral inputs to CNNs. In support of this model, brain slice studies have identified an NMDA receptor-dependent form of long-term potentiation of excitatory inputs to CNNs that is triggered upon release of hyperpolarization (Pugh and Raman, 2006). Our findings indicate that training results in increases in presumed mossy fiber innervation of areas within the cerebellar nuclei targeted by modulated PNs (Figure 5), consistent with a strengthening of tone-related mossy fiber inputs. The results demonstrate that release from hyperpolarization caused by the

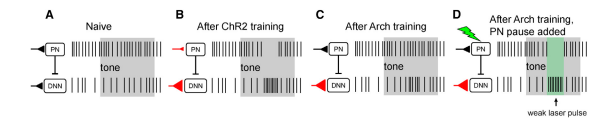


Figure 8. Proposed Circuit Basis of ChR2- and Arch-Driven Learning

(A–D) Schematic of proposed model for associative learning in ChR2 and Arch animals.

(A) Before learning, the tone (gray box) does not drive activity changes in CNNs.

(B) After learning, PNs in ChR2 animals respond with tone-evoked pauses that disinhibit CNNs. Changes in synaptic strength of PN and CNN inputs (red triangles) are hypothesized to underlie learned changes in excitability.

(C) After Arch training, learning results only in increased excitability of CNNs to the learned response.

(D) Weak Arch activation (green box) timed to the peak of the forelimb movement mimics disinhibition of CNNs, thereby producing stronger CNN-driven motor output.

simultaneous pause in firing of a population of PNs can serve as the teaching signal driving such plasticity.

Ensuring precise timing of movements is a hallmark of cerebellar function, as demonstrated for classically conditioned eyeblinks (Ohyama and Mauk, 2001; Ohyama et al., 2006; Perrett et al., 1993). Pharmacological inactivation of the cerebellar cortex after eyeblink training does not abolish LRs, but leads to rapid eyeblinks with time courses that are no longer in register with the training interval. The Arch-induced learning described here shows a similar profile (Figure 6), consistent with learning under these circumstances being confined to the cerebellar nucleus (McCormick and Thompson, 1984).

A unifying hypothesis accounting for optogenetic and normal forms of learning is presented in Figure 8. We posit that pairing ChR2 excitation of PNs with sensory stimuli triggers changes in synapses encoding those sensory stimuli at PNs and CNNs, leading to sensory-evoked pauses in PNs and bursts in CNNs, and robust, well-timed learned movements. In contrast, pairing Arch stimulation with sensory stimuli leads only to changes in inputs to CNNs (compare Figures 8B and 8C). We hypothesize that less robust and less well-timed motor-learned movements in Arch-trained mice occur because sensory-evoked CNN bursts do not occur coincidently with PN disinhibition (Figure 8C). Supporting this interpretation, we demonstrate that in Arch-trained mice, imposition of pauses in upstream PNs (Figure 8D) facilitates movements elicited by the sensory conditioning stimulus (Figure 7). Taken together, these results provide further evidence that cerebellar-mediated learning recruits plastic changes at multiple sites within the circuit.

Functional Implications

By generating artificial motor memories through optogenetic training, we show that the effects of PN pauses on movement are augmented by learning, providing support for models of cerebellar learning that hypothesize learning-related circuit changes in the cerebellar nucleus (Miles and Lisberger, 1981; Perrett et al., 1993). Our results further show that disinhibition of CNNs is sufficient to drive associative memory formation. The fact that PN pauses can instruct memory formation provides proof of concept support for theories of cerebellar memory formation that propose that learning-related changes in the cere-

bellar cortex precede, and may be necessary for, structural plasticity and consolidation in the cerebellar nucleus (Cooke et al., 2004; Kassardjian et al., 2005; Ohyama et al., 2006).

EXPERIMENTAL PROCEDURES

Animals

All animal procedures were performed in accordance with NIH standards and were approved by the University of California, Los Angeles Institutional Animal Care and Use Committee. Male and female mice homozygous for L7-Cre (B6.129-Tg(Pcp2-cre)2Mpin/J, The Jackson Laboratory) were crossed with either an animal homozygous for ChR2-eYFP (Ai32, B6;129S-*Gt*(*ROSA*)26Sor^{tm32(CAG-COP4+H134R/EYFP)Hze/J, The Jackson Laboratory) or Arch-eGFP (Ai35, B6;129S-*Gt*(*ROSA*)26Sor^{tm35.1(CAG-aop3/GFP)Hze/ J, The Jackson Laboratory). Subsets of animals at the end of behavioral experiments were perfused (4% paraformaldehyde) and their brains removed. Fluorescent images of the whole cerebellum were obtained at 14× using a Zeiss (Stereo Discovery V12) dissecting microscope and camera (Axiocam MRm) in order to examine and document the sites of chronic fiber placement in each mouse (Figure S1).}}

In Vitro Electrophysiology

Adult mice 23–34 g (L7-Cre/Arch or ChR2) in weight were anesthetized and decapitated. Parasagittal cerebellar slices (300 μ m) were cut in an ice-cold (4°C), low-sodium cutting solution using a vibratome (Leica VT1000). Slices were incubated for ~30 min at 35°C and allowed to sit at room temperature before electrophysiological recordings at ~34°C. Cutting and recording media were bubbled with 95% O₂ and 5% CO₂. The low-sodium cutting solution consisted of (in mM) 82.7 NaCl, 2.4 KCl, 1.4 NaH₂PO₄, 0.5 CaCl₂, 6.8 MgCl₂, 23.8 NaHCO₃, 65 sucrose, and 23.7 dextrose, and the recording solution consisted of (in mM) 119 NaCl, 2.5 KCl, 1.25 NaH₂PO₄, 2 CaCl₂, 1 MgCl₂, 26 NaHCO₃, and 25 dextrose.

Cells were visualized using an upright microscope (Zeiss Axioskop II) with a 40× water immersion lens using infrared differential interference contrast microscopy. Optical activation of ChR2 or Arch was achieved using an LED light source (532 nM, THORLABS) projected through the epifluorescence pathway of the microscope. Pulses of light were either triggered directly by TTL signals from pClamp to the LED controller or through a signal generator (Master 8, A.M.P.I.) triggered by pClamp. For all graphs, optical stimulation was elicited at t = 0 ms. Data were acquired using pClamp9 (Molecular Devices) at 50 kHz for intracellular recordings and 5–10 kHz for extracellular recordings. A MultiClamp 700B was used to record electrophysiological signals. In voltage clamp, the pipette and cellular capacitance ($\geq 80\%$) were compensated for using onboard circuitry. The pipette solution for PN voltage and current clamp recordings contained (in mM) 126 KMSO₃, 10 KCl, 4 NaCl, 10 HEPES, 0.5 EGTA, 14 Tris-phosphocreatine, 2 MgATP, and 0.4 NaGTP. Experiments

were analyzed using custom macros written for Igor Pro (Taro Tools, Dr. Taro Ishikawa, http://sites.google.com/site/tarotoolsregister).

Surgical Procedures

All surgical procedures were performed under isoflurane (1%) anesthesia, and performed at least 2 days prior to either in vivo electrophysiological recording or behavioral manipulation. Animals were placed into a stereotaxic device, and custom-made head bars (Fab 2 Order) were glued to the skull using Vetbond (3M) and dental cement (Bosworth). For in vivo electrophysiological recordings large craniotomies were made over the medial or left cerebellum. The exposed site was then filled over with a silicon-based elastomer (Kwik-Cast, World Precision Instruments) that was easily removed just prior to recording. For behavioral experiments, chronically implanted optic fiber cannulas (Doric Lenses), dipped in Dil (Sigma), were stereotaxically positioned (RC, 6.25 mm; ML, 1.9 mm; DV, 2 mm) into the brain through small craniotomies, and fixed into place using Metabond (Parkell). Whole-mount fluorescence visualization (see Figure S1) allowed post hoc localization of the fiber.

In Vivo Electrophysiology

Head-fixed animals were allowed to move in place freely on a spinning disk (Ware Flying Saucer). Optrodes constructed of fiber-optic cannulas (Doric Lenses) glued ~400 μm behind 1–5 M Ω Parylene-C-insulated tungsten electrodes (A-M Systems) were inserted into craniotomies and vertically driven into the cerebellum using a micromanipulator (Sutter). Electrical signals were recorded with an Axopatch 200B amplifier (V-clamp mode), and data were acquired in pClamp9 (50 kHz). A TTL-controlled, 100-W diode-pumped solid-state laser (532 nm, Opto Engine) was coupled through a patch cable (Doric Lenses) to the optrode to deliver brief pulses of light. Spike detection and data analysis were performed in Igor Pro 6 using custom macros (adapted from Taro Tools).

Behavior

All animals were habituated to the behavioral setup for at least 4 days prior to training. Animals were head-fixed, but allowed to move freely on a spinning disk (Ware). A TTL-controlled, 100-W, 532-nm diode laser (CNI Laser, Opto Engine), launched into a patch cable and connected to the fiber-optic cannula, was used to activate either ChR2 or Arch within the forelimb region of the cerebellum for the indicated durations determined by TTL control. Power output per unit area, measured regularly from a pristine fiber segment identical to that implanted, ranged from 320 to 1,910 mW/mm². These values are calculated by dividing total power (10–60 mW) by the cross-sectional area of the 200-µm diameter optical fiber used in our study. The values used here compare well to values of 142–16,000 mW/mm² reported by Heiney et al. (2014) for the 20- to 30-µm diameter fibers used in their study.

Optical stimulation was paired with a tone coordinated using custom routines written in LabView and controlled via TTL pulses from a NI-DAQ board (National Instruments). Epochs of high-speed video (200 f/s) were also synchronized via TTL and obtained with a Giga-E camera (Allied). Kinematic measurements were made offline using Custom LabView routines that allowed for the tracking of an IR reflective button (MoCap Solutions) adhered to the mouse's wrist. Data were further analyzed using custom macros in Igor Pro 6 allowing infrequent tracking artifacts to be excised. A subset of the kinematic data were also analyzed by MTrackJ in ImageJ, and the results were identical to the automated LabView routine. All speed-versus-time traces were smoothed by a binomial function with a factor of 3. All error bars or shaded errors in figures represent SEM.

Custom Igor procedures were used to automatically sort individual trials for rejection or analysis. Rejected trials included those in which movement speed exceeded 0.04 m/s during a 500-ms epoch prior to tone onset (baseline period) or trials in which movement speed exceeded 0.04 m/s within 50 ms after tone onset (startle period). On average this resulted in variable rejection rates of 35–70 trials per mouse per day. In trials with tone and laser stimuli, LRs were judged to occur when movement speed exceeded 0.04 m/s in the time epoch from 50 ms after tone onset until laser onset; in tone-alone trials LRs could occur from 50 ms following tone onset until tone offset, which co-terminated with the laser pulse offset. To determine percent LR, the number of LR trials was divided by total number of analyzed trials (excluding rejected

trials). We chose the term "learned response," rather than the more conventionally used term "conditioned response," to reflect the fact that LRs in this study are generated by optogenetic stimuli, rather than by behavioral error signals, and because LRs in ChR2- and Arch-trained animals are likely to result from distinct circuit mechanisms.

PN Tract Tracing

Dextran-conjugated Alexa 488 (210 nL, 5 mM) was stereotaxically injected into the cerebellar cortex of wild-type mice at the same coordinates used for fiber implants. After allowing retrograde transport of the dye for 7 days, mice were anesthetized (pentobarbital) and then perfused with a 4% paraformaldehyde phosphate-buffered saline (freshly diluted from 32% stock, Electron Microscopy Sciences). Brains were subsequently removed and incubated in a phosphate-buffered solution containing 30% sucrose for up to 2 days. Brains were then embedded in optimal cutting temperature solution (TissueTech) at -80°C. Sections, cut in a cryostat at 60 µm, were immunostained using mouse α-NeuN monoclonal antibody (1:500, Millipore) and goat α-mouse Alexa 568 secondary antibody (1:300, Life Technologies). Confocal images were obtained at 10 and 20× magnification (0.3 and 0.8 NA objectives, respectively) using a Zeiss 710 laser-scanning confocal microscope. Raw images were processed and analyzed using ImageJ (NIH). Quantification of dye-filled axons was performed by first thresholding images of sagittal or coronal cerebellar sections and then calculating the percent immunofluorescence in each nucleus (Lat = $0.013\% \pm 0.007\%$, n = 7 sections: IntA = $20.5\% \pm 5.93\%$, n = 17: IntP = 6.87% ± 1.85%, n = 17; MedDL = 38.8% ± 5.38%, n = 17; Med = $4.81\% \pm 1.45\%$, n = 16; data from two animals). Each section was also visually examined for the presence of dye-filled PN axons (Lat = 0/12 sections, IntA = 12/12, IntP = 13/17, MedDL = 16/16, Med = 14/17; n = two animals). Any labeled tubular structures with thicknesses of more than 3 µm were presumed to be endothelial cells composing blood vessels, which were found throughout the sections and thus excluded from analysis (see Figure S7).

Learning-Induced Structural Plasticity in the Cerebellar Nuclei

L7-Arch mice were habituated on the behavioral platform for at least 4 days. Cage siblings were split into paired (100 pairs of 250-ms, 2-kHz tone co-terminating with 75-ms laser stimulation, randomized inter-trial interval between 20 and 40 s) and unpaired (100 tone-alone/laser-alone trials, randomized intertrial interval between 10 and 20 s) training groups. For the unpaired training, tone and laser presentation was performed in a pseudorandomized order. After training for 4 consecutive days, the mice were perfused with 4% paraformaldehyde and prepared for cryosectioning as described above. Parasagittal sections (30 µm) were then cut and stained with mouse α-VGluT2 monoclonal antibody (1:300, Millipore) and goat a-mouse Alexa 568 secondary antibody (1:300). Images of the cerebellum were first obtained using a Zeiss 710 confocal microscope with a 10× objective to identify the cerebellar nuclei. Each identified region of the cerebellar nuclei was then captured as continuous stacks of confocal images (0.64 µm thick) using a 63× objective (Zeiss, 1.4 NA) and custom acquisition software (Zeiss). After image acquisition, every fifth image in the z stack was isolated to prevent recounting the same puncta. The VGluT2+ puncta were quantified by first thresholding the image to isolate immunopositive signals. The particle count tool in ImageJ (NIH) was then used to count discrete puncta in each identified cerebellar nucleus.

SUPPLEMENTAL INFORMATION

Supplemental Information includes seven figures, one table, and four movies and can be found with this article online at http://dx.doi.org/10.1016/j. neuron.2015.03.010.

AUTHOR CONTRIBUTIONS

K.H.L., P.J.M., A.M.B.R., K.Y.C., and T.S.O. designed experiments. K.H.L. performed the initial experiment observing forelimb movement to launch the project and the surgery and behavioral learning experiments in Figures 3, 4, and 6. P.J.M. performed in vivo and in vitro recordings, made behavioral kinematic measurements with S.A.J., and generated the histological

reconstructions with A.M.B.R. and K.Y.C. K.Y.C. conducted behavioral training on paired and unpaired mice as well as performed the histology and analysis of data in Figures 5 and S7. A.M.B.R. performed the augmentation experiment in Figure 7, wrote Igor routines used to analyze data in Figures 3–6, and analyzed the data in Figures 7, S5, and S6. R.E.S. designed the custom software for acquisition and analysis of the behavioral data. P.J.M. and T.S.O. wrote the manuscript with help from K.H.L., K.Y.C., and A.M.B.R. Given the equal contributions made by K.H.L. and P.J.M., the first author position was determined by a coin flip.

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